DISTRICT OF COLUMBIA \sim DEPARTMENT OF HEALTH \sim ADAP

Sofosbuvir tablet (Solvadi®)

PRIOR AUTHORIZATION PROGRAM Request Form – Initial Request

CLIENT'S NAME: _____ ADAP ID: _____

CLIENT	'S DATE OF BIRTH_	AI	DAP Pharmacy		
(HC\		,	nucleotide analog inhibito lable as a 400mg film-coa	•	
Solv requ	•	ior approval for cover	rage. Allow up to 96 hou	rs for completion of	
			necessity of necessity (knowledgement and con		
Sofosk of a co The ef infection for live (HCV/	ombination antivir fectiveness of so on, including pation	al treatment regimen. fosbuvir was establishe ents with hepatocellular d those with hepatitis C on.	onic hepatitis C (CHC) info d in patients with HCV ge carcinoma meeting Milan virus/human immunodefio	notype 1, 2, 3 or 4 criteria (those waiting	
Drug	ulo Heatillelit K	Dose	Route	Frequency	
			1.00.00	. requestey	
Criteri	ia for use:				
Please	complete and chec	k all that apply:			
1.		ase specialist or gastroente	of HIV/hepatitis C infection erologist.	n, or in consultation with	
2.	Client does have adherence issues with antiretroviral or other medications. YES NO				
3.		receiving or recently rece	eived amiodarone.		
4.	Client's has confirmed clinical diagnosis of Hepatitis C, genotype YES \square NO \square				
5.	Client is not pregnant. YES N		ome pregnant and/or female p	partner of a male patient is	
6.	Client does have	decompensated liver disea	ise.		

YES □ NO) 🗆						
7. Client has cirrhosis	_						
YES □ NO) 🗆						
8. Client will not be treated with medications that are not recommended for use with or							
contraindicated wit	h sofosbuvir						
YES □ NO) 🗆						
9. Client has a FibroSure score of							
	c or biopsy proven sco	re of Date:					
	sitive hepatitis C viral load take						
	start date of Solvadi® is						
	duration of CHC treatment is _	weeks.					
P							
Recommended dosage	and administration: The rec	commended dose is one 400mg					
tablet orally once a day with	or without food. Sofosbuvir should	be administered in combination with					
ribavirin or in combination with peginterferon and ribavirin to treat chronic hepatitis C in adults. The							
following table describes the duration of therapy for adults with sofosbuvir combination therapy in							
-	r in patients co-infected with CHC/F						
patients injected with One o	In patients co-infected with orion						
Onfortherin Topoter and	Deniment and Denetions has also	Detient Oberesteristics (Defenses of	. ()				
		on Patient Characteristics (Reference On	ily)				
Genotype	Treatment	Duration					
Genotype 1 or 4 CHC	Sofosbuvir plus peginterferon alfa plus ribavirin	12 weeks					
Genotype 2 CHC	Sofosbuvir plus ribavirin	12 weeks					
Genotype 3 CHC	Sofosbuvir plus ribavirin	24 weeks					
The dose of ribavirin is weight-	based, as determined by the manu	facturer's guidelines. For patients with					
genotype 1 CHC, an alternative	e for patients ineligible for interferor	therapy is the administration of					
sofosbuvir plus ribavirin for 24	weeks.						
The recommendation for the us	se of sofosbuvir in patients with her	patocellular carcinoma awaiting liver					
transplantation is combination	therapy with ribavirin for up to 48 w	eeks or until the time of the transplant					
(whichever comes first).							
Dhysisian/s signature.		Data					
Physician's signature:		Date:					
Physician's Name (Print):_	Phor	ne #:Fax #:					
Fax Completed Form to Cli	nical Pharmacy Associates: Fax	: 1 (888) 971-7229					
•	•						
FIIUIIE. 1 (000) /43-0434 E	xt 150 Attention: Prior Approv	ai Fiografii					
Ammanal: VEC = NO	Data	044:					
		s Office use only					
Reason for denial							

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